1 The Impact of Antimalarial Resistance on the Genetic Structure of *Plasmodium falciparum* in the 2 DRC

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40 ABSTRACT

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42 The Democratic Republic of the Congo (DRC) harbors 11% of global malaria cases, yet little is known 43 about the spatial and genetic structure of the parasite population in that country. We sequenced 2537 44 Plasmodium falciparum infections, including a nationally representative population sample from DRC 45 and samples from surrounding countries, using molecular inversion probes - a novel high-throughput 46 genotyping tool. We identified an east-west divide in haplotypes known to confer resistance to 47 chloroquine and sulfadoxine-pyrimethamine. Furthermore, we identified highly related parasites over 48 large geographic distances, indicative of gene flow and migration. Our results were consistent with a 49 background of isolation by distance combined with the effects of selection for antimalarial drug 50 resistance. This study provides a high-resolution view of parasite genetic structure across a large 51 country in Africa and provides a baseline to study how implementation programs may impact parasite 52 populations.

53 BACKGROUND

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55 Malaria remains one of the largest global public health challenges, with an estimated 219 million cases 56 worldwide in 2017¹. Despite decades of scale-up in control, there has been a recent resurgence, 57 particularly in high transmission countries in sub-Saharan Africa¹. In addition, the emergence of 58 antimalarial resistance poses a major threat to current control and elimination efforts worldwide, and 59 new tools are needed to quantify the changing landscape of drug resistance on timescales relevant to 60 malaria control programmes. Genomics has emerged as a useful method for better understanding 61 parasite populations that can be leveraged to support the design of effective interventions against a 62 continually evolving parasite.

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Data from genomic studies provides information that is complementary to epidemiological data², and 64 65 can help to answer several key questions, including how parasites are transmitted, how drug resistance 66 spreads, and how malaria control efforts impact the diversity of the parasite population. However, to 67 date, efforts to use genomics to inform malaria control efforts have suffered from three major 68 limitations. First, much of the work has been conducted in low transmission regions, such as Asia and 69 transmission fringe regions of Africa, leaving it unclear how useful information can be gathered in the 70 highest transmission settings. Some of these high burden regions have experienced increasing malaria 71 prevalence in recent years and are now the center of strategic plans for control efforts^{3,4}. Second, most 72 genomic studies in Africa have relied upon convenience sampling from a few sites usually collected for 73 other purposes, rather than population-representative samples. Lastly, studies have either relied on 74 relatively few genetic markers, providing limited insight into the complete genome, or on expensive 75 whole genome sequencing, limiting the number of samples studied. Overcoming these limitations is 76 essential for genomics to have broader impacts on malaria control.

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78 Within Africa, parasite populations have been shown to vary significantly between East and West, as demonstrated by their distinct antimalarial drug susceptibilities and population genetics^{5,6}. However, few 79 80 genomic studies have incorporated samples from central Africa, limiting our understanding of the 81 connectivity of parasite populations across the continent. The Democratic Republic of the Congo (DRC) 82 is the largest malaria-endemic country in Africa, borders nine countries and harbors approximately 11% of global *P. falciparum* malaria cases¹. The DRC harbors a large, understudied parasite population that 83 84 likely serves as a bridge between African parasite populations. Limited previous work has shown that 85 the DRC represents a watershed between East and West African drug resistant parasite populations for sulfadoxine-pyrimethamine and chloroquine resistance⁷⁻⁹. More recently, parasite population structuring 86

due to mutations at these and other loci associated with antimalarial resistance has been confirmed
within the DRC¹⁰. However, studies focusing on hypervariable surface antigen diversity or neutral
microsatellites have been unable to detect significant structure in the parasite population^{10,11}, likely due
to a lack of high-quality genome-wide signal. A better understanding of parasite populations and the
spread of antimalarial resistance in the DRC will allow for the design of more effective interventions
accounting for evolutionary forces.

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94 To address this knowledge gap, we leveraged a recent advance in malaria genomics, high-throughput 95 molecular inversion probe (MIP) capture and sequencing, to characterize and map parasite population 96 structure and antimalarial resistance profiles in the DRC and to define the connections of parasites within the DRC to East and West African parasite populations¹². This approach provides a cost-97 98 effective and scalable method of genome interrogation, without the expense or informatic complexities 99 of whole genome sequencing. We previously employed MIPs to comprehensively genotype known antimalarial resistance genes in several hundred samples from the DRC¹⁰. Here, we introduce an 100 101 expanded MIP panel targeted at 1834 single nucleotide polymorphisms (SNPs) distributed throughout 102 the *P. falciparum* genome, and designed to quantify differentiation and relatedness between samples. 103 Using this panel of genome-wide SNP MIPs, in combination with the previous drug resistance MIP 104 panel, we evaluated the parasite population diversity in 2537 parasite isolates from the DRC and 105 surrounding countries in East and West Africa. We used this information to quantify relatedness of and 106 gene-flow between parasites over large geographic scales and to assess the origins of antimalarial 107 resistance mutations.

108 **RESULTS**

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110 Sample guality and filtering: We obtained 2537 samples collected in 2013-2015 from the DRC and 111 surrounding countries (DRC=2039, Ghana=194, Tanzania=120, Uganda=63, Zambia=121). All 112 samples were sequenced using two separate MIP panels: a genome-wide panel designed to capture 113 overall levels of differentiation and relatedness, and a drug resistance panel designed to target 114 polymorphic sites known to be associated with antimalarial resistance¹⁰. The genome-wide panel 115 included 739 ostensibly geographically informative SNPs, chosen on the basis of high differentiation 116 (F_{ST}) between surrounding African countries in publicly available genomic sequences made available 117 by the Pf3K project (see **Supplemental Text 1** and **Supplemental Table 1**), and 1151 putatively 118 neutral SNPs distributed throughout the genome, with an overlap of 56 SNPs that were both neutral 119 and geographically informative. The drug resistance panel included SNPs in known and putative drug resistance genes and has been described elsewhere ¹⁰. The median number of unique molecular 120 121 identifiers (UMIs) per MIP was 31 (range: 1-8,490) for the genome-wide panel, and 10 (range: 1-122 32,511) for the drug resistance panel. Complete UMI depth distributions are shown in Supplemental 123 Figure 1. After filtering for samples and loci with sufficient UMI coverage, we were left with 1382 124 samples and 1079 loci from the genome-wide panel, and 674 samples and 1000 loci from the drug 125 resistance panel, with an overlap of 452 samples between both panels. In addition to these samples, 126 114 controls consisting of known mixtures were sequenced and used to assess the accuracy of allele 127 calls and frequencies. Expected versus measured allele frequencies for each SNP, calculated from 128 these controls, are shown in Supplemental Figure 2.

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130 **Complexity of infection:** Initial analyses focused on the genome-wide MIP panel only. Complexity of 131 infection (COI) for each sample was estimated using THE REAL McCOIL¹³ (Supplemental Figure 3). 132 The mean COI was estimated at 2.2 (range 1 - 8) for the study as a whole. We observed significant 133 differences in COI between countries (Ghana: 1.55 (non-parametric bootstrap 95% CI: 1.39 - 1.73), 134 DRC: 2.23 (2.15 - 2.31), Tanzania: 2.17 (1.83 - 2.51), Zambia: 2.68 (2.39 - 3.00), Uganda 2.18 (1.87 -135 2.51), and within the DRC we observed a statistically significant relationship between COI and P. 136 falciparum prevalence by microscopy at both the province and cluster levels (Supplemental Figure 4), 137 with higher COIs observed at higher prevalences.

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Population structure: We explored population structure through principal component analysis (PCA)
evaluated on within-sample allele frequencies at all 1079 genome-wide loci. We found the same
separation between East and West Africa described in previous studies (Figure 1) as well as finer

142 structure between regions within East Africa. DRC samples comprised a continuum between the East



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Figure 1 The first two (a) and three (b) principal components calculated from within-sample allele frequencies using the 147 genome-wide MIP panel. Colors indicate country of origin of each sample. 148

149 The relative contribution of each locus to each principal component was quantified through normalized 150 loading values. Relative contributions to the first four principal components are shown in Figure 2. After 151 the fourth principal component the percent variance explained by subsequent components plateaued 152 (Supplemental Figure 5). For principal component 1 (PC1) large contributions came from loci 153 distributed throughout the genome, and a relatively larger contribution (65.2%) came from putatively 154 geographically informative SNPs (non-parametric bootstrap, p<0.001). In contrast, contributions to PC2 155 were concentrated in a region on chromosome seven in close proximity to P. falciparum chloroquine 156 resistance transporter (*pfcrt*), a known drug resistance locus, suggesting that resistance to chloroguine 157 or amodiaguine may be driving differentiation along this secondary axis. For PC3, locus contributions were concentrated in three genic regions: PF3D7 0215300 (8.5%), PF3D7 0220300 (5.0%), and 158 159 PF3D7 1127000 (4.3%). The first and largest of these encodes an acyl-CoA synthetase and is part of a diverse gene family known to undergo extensive gene conversion and recombination¹⁴. For PC4 we 160 161 observed a region of high locus contribution on chromosome eight in close proximity to the known 162 antifolate drug resistance gene dihydropteroate synthase (*dhps*). Combined, these results suggest that 163 geography and drug resistance are both contributors to the observed population structure.

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- 165 The relationship between the PCA results and the spatial distribution of parasites was explored by
- 166 plotting raw principal component values against the geographic location of samples (Figure 3a-3d). For
- 167 PC1 this revealed a complex pattern of spatial variation, containing both north-south and east-west
- 168 clines. For PC2 and PC4 the maps essentially recapitulate the known geographic distribution of *pfcrt*
- and *dhps* resistance mutations, respectively (Figure 3e-3f). For PC3 the map indicates some east-west
- 170 spatial structuring that is not explained by known markers of antimalarial resistance and warrants
- 171 further investigation.



172

173Figure 2 The relative contribution (%) of each locus to the first four principal components. Chromosomes are plotted in order,174separated by vertical white gridlines. Point colors indicate sites that were chosen in the design based on F_{ST} values to be175geographically informative (blue) or not (red).



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180 Identity by Descent: The relatedness of all pairs of samples was explored through pairwise identity by 181 descent (IBD), estimated using a maximum likelihood approach. IBD has advantages over simpler 182 statistics like identity by state (IBS) in that it takes account of allele frequency distributions, and so 183 provides an objective measure of relatedness that can be compared between studies¹⁵. The overall 184 distribution of pairwise IBD was found to be heavy-tailed, consisting of a large body of weakly related 185 samples and a tail of very highly related samples (Figure 4).

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Figure 4 A histogram of pairwise identity by descent (IBD) between all samples, estimated by maximum likelihood. Inset shows the heavy tail of the distribution, with some pairs of samples having IBD > 0.9.

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191 Mean IBD was significantly higher within clusters compared to between clusters (0.06 vs. 0.02, two-192 sample t-test, p<0.001). When plotted against geographic separation there was a clear fall-off of IBD 193 with distance (Figure 5a), consistent with the classical pattern expected under isolation-by-194 distance^{16,17}. Focussing on the tail of highly related samples, which includes the major strain in complex 195 infections, there were 12 sample pairs with a relatedness greater than IBD=0.9. Comparison of raw 196 allele frequency distributions confirmed that these were likely clones (Supplemental Figure 6). These 197 highly related pairs were found more often within the same cluster than in different clusters (7 vs. 5 198 respectively, chi-squared test, p<0.001), suggesting the presence of local clonal transmission chains. 199 The five between-cluster highly related pairs (Figure 5b) were spread over large geographic distances 200 (281-1331 km), far beyond the normal expected scale of the breakdown in genetic relatedness (Figure 201 5a), suggesting recent long distance migration.

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Figure 5 Panel (a) shows the mean IBD between clusters, binned by the spatial distance between clusters. Vertical lines show
 95% confidence intervals. Panel (b) shows the spatial distribution of highly related (IBD>0.9) parasite pairs. Black areas
 indicate major water bodies, including the Congo River.

208 Prevalence of markers of resistance: Based on previous findings of an east-west divide in molecular markers of antimalarial resistance in the DRC^{8,9}, all samples in the DRC were divided by 209 210 aeographically-weighted K-means clustering into two populations (**Supplemental Figure 7**). The 211 prevalence of every mutation identified by the drug resistance MIP panel was calculated in eastern and 212 western DRC, as well as at the country level. **Table 1** gives a summary of all mutations that reached a 213 prevalence >5% in any geographic unit, and a complete list of all identified mutations along with their 214 prevalence is given in **Supplemental Table 2**. Note that in the *dhps* mutation **G**437A the reference is 215 resistant, hence this is re-coded as A437G and prevalence values indicate the prevalence of the 216 reference allele. Estimated prevalences of these alleles in the DRC as a whole were broadly similar to previously published estimates¹⁰. However, we did identify several polymorphisms in known and 217 218 putative resistance genes not previously reported in the DRC, including kelch K189T and pfatp6 N569 \underline{K} , both of which have been described at appreciable frequencies elsewhere in Africa^{18–20}. 219 220 221 Geographic distribution of haplotypes: Previous studies have demonstrated that mutations associated with antimalarial resistance are clustered into east-west groupings within DRC^{8,10}. Focusing 222 223 on the 107 samples from DRC that were identified as monoclonal from The REAL McCOIL analysis, we 224 explored the joint distribution of all combinations of mutant haplotypes in both the *dhps* and *crt* genes. 225 Raw combinations of mutations were visualized using the UpSet package in R²¹, and the spatial 226 distribution of haplotypes in the DRC was explored by plotting these same mutant combinations against

their corresponding DHS cluster locations (**Figure 6**). Our results for *dhps* recapitulate those found

previously, showing a clear east-west divide with the K540<u>E</u> and A581<u>G</u> mutants concentrated in the

east, and S436<u>A</u> and A437<u>G</u> concentrated in the west. For *crt* we also find evidence of an east-west

divide, with haplotypes containing N326<u>S</u> and F325<u>C</u> concentrated in the east and those containing

- 231 I356<u>T</u> concentrated in the west.
- 232

				prevalence						
gene	chromosome	position	mutation name	overall	DRC	DRC West	DRC East	Ghana	Uganda	Zambia
atp6	chr1	267007	1723V	1.1	0.3	0.7	0.0	4.2	7.3	0.0
atp6	chr1	267257	G639D	2.0	1.8	2.9	1.0	0.0	7.3	0.0
atp6	chr1	267467	N569K	24.1	21.9	18.8	24.0	16.7	41.5	28.9
atp6	chr1	267882	E431K	15.3	17.0	18.8	15.7	16.7	9.8	6.7
atp6	chr1	267970	L402V	7.1	8.2	10.1	6.9	12.5	0.0	2.2
dhfr-ts	chr4	748239	N51I	83.0	79.5	81.2	78.4	75.0	100.0	97.8
dhfr-ts	chr4	748262	C59R	71.2	63.2	63.0	63.2	95.8	95.1	97.8
dhfr-ts	chr4	748410	S108N	97.8	97.1	97.1	97.1	100.0	100.0	100.0
dhfr-ts	chr4	748577	I164L	3.1	0.6	0.0	1.0	0.0	29.3	0.0
mdr1	chr5	958145	N86Y	12.4	14.3	18.8	11.3	16.7	7.3	0.0
mdr1	chr5	958440	Y184F	37.4	36.5	39.9	34.3	58.3	31.7	37.8
mdr1	chr5	958484	T199S	1.3	0.0	0.0	0.0	0.0	14.6	0.0
mdr1	chr5	958584	S232Y	2.7	3.5	5.1	2.5	0.0	0.0	0.0
mdr1	chr5	961625	D1246Y	4.4	2.9	3.6	2.5	0.0	24.4	0.0
crt	chr7	403620	M74I	30.3	28.7	37.7	22.5	16.7	85.4	0.0
crt	chr7	403621	N75E	30.3	28.7	37.7	22.5	16.7	85.4	0.0
crt	chr7	403625	К76Т	30.3	28.7	37.7	22.5	16.7	85.4	0.0
crt	chr7	404407	A220S	28.1	24.6	31.9	19.6	8.3	100.0	0.0
crt	chr7	405600	I356T	7.1	9.4	21.0	1.5	0.0	0.0	0.0
dhps	chr8	549681	S436A	15.0	17.3	28.3	9.8	37.5	0.0	0.0
dhps	chr8	549685	A437G	73.2	67.3	72.5	63.7	95.8	100.0	82.2
dhps	chr8	549993	K540E	25.4	17.0	9.4	22.1	0.0	85.4	48.9
dhps	chr8	550117	A581G	8.2	6.1	2.2	8.8	0.0	34.1	4.4
k13	chr13	1726431	K189T	14.8	14.9	18.8	12.3	54.2	0.0	6.7
mdr2	chr14	1956202	1492V	23.2	21.3	22.5	20.6	20.8	31.7	31.1
mdr2	chr14	1956408	F423Y	31.4	30.1	28.3	31.4	29.2	36.6	37.8

233

Table 1 Prevalence (%) of mutations identified by the drug resistance MIP panel. Includes all mutations that reached a
 prevalence >5% in any given geographic unit.

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237 Selective sweep and haplotype analysis: Using the drug resistance MIPs and genome-wide SNP 238 MIPs combined, the extended haplotypes of the monoclonal infections were determined for 200kb 239 upstream and downstream of each putative drug resistance allele that had at least 5% overall 240 prevalence in the DRC. The CVIET haplotype within the *crt* gene showed a signal of positive selection. 241 with longer haplotype blocks in western DRC as compared to eastern DRC (**Figure 7**; p'XP-EHH_D < 242 0.05). In the east, patterns of haplotype homozygosity are consistent with positive selection for the 243 derived I356T haplotype (Supplemental Figure 8), although a XP-EHH_D statistic could not be 244 calculated for this locus because the derived haplotype was absent in western DRC, supporting the 245 geographic localization of the I356T mutation in the east (Figure 6).



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Figure 6 The spatial distribution of all combinations of mutant haplotypes for *dhps* and *crt* from the monoclonal DRC samples. Panels (a) and (c) are UpSet plots showing the number of times each combination of mutations was seen for *dhps* and *crt*, respectively. Panels (b) and (d) show these same haplotypes on a map of DRC. Colours correspond horizontally between panels, i.e. between (a) and (b), and between (c) and (d), with the exception of wild-type haplotypes (grey) which are not shown in panels (b) and (d).

253 Mutations in *dhps* were more difficult to interpret. This gene has undergone multiple selective sweeps 254 associated with increasing drug resistance. The most recently introduced mutation into the DRC, dhps 255 A581G, showed relatively conserved local haplotypes around the mutation in both eastern and western 256 DRC (Supplemental Figure 9). Extended haplotypes around the other mutations (Supplemental 257 Figures 10 and 11) are inconsistent with a classical hard sweep, perhaps due to selection on multiple 258 independent haplotypes or to interference between A581G and other linked alleles. Finally, we did not 259 detect any strong signals of differing patterns of recent positive selection between the eastern and 260 western DRC among the *dhfr* and *mdr*² genes (Supplemental Table 3, Supplementary Figure 12). 261



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Figure 7. EHH and Bifurcation Plots for *pfcrt* K76T from the monoclonal samples with no missing genotype data. Panels (a) and (b) display EHH curves 200 kb upstream and downstream from the K76T core SNP in centimorgans among the samples from the eastern DRC and western DRC. Panels (c) and (d) show haplotype bifurcation plots with respect to the core allele ancestry and the eastern DRC and western DRC for a subsetted region. Position is considered in kilobases, and segregating sites for each haplotype are displayed at the nodes. Overall, there is strong evidence for recent positive selection of the *pfcrt* CV<u>IET</u> haplotype in the west that is mitigated in the east.

269 DISCUSSION

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271 Here we provide the first large-scale, robustly sampled study of *falciparum* malaria in central Africa 272 using MIP capture and sequencing, a novel high-throughput genotyping approach that is appropriate for 273 large population based surveys. Using a panel of probes designed to detect genome-wide SNPs, 274 combined with a second panel targeting drug resistance genes, we were able to show that the parasite 275 population in the DRC contains a signal of differentiation by geographic separation, consistent with the 276 classical pattern of isolation by distance. This background population structure is overlaid with the clear 277 impacts of drug resistance mutations, which cause distinct structure between East and West African 278 parasite populations. Additionally, the use of relatively dense genome-wide SNPs allowed us to carry 279 out relatedness analysis, revealing a handful of cases where human hosts separated by many 280 hundreds of kilometers were infected by essentially identical clones. Given the rapid breakdown of 281 distinct genotypes by recombination in high transmission areas, it is highly likely that these events 282 represent relatively recent infection and migration events. With this in mind, it is interesting to note that 283 pairwise links of high relatedness tend to fall along the Congo River, an important route of 284 transportation in DRC. Lastly, the combination of the two MIP panels allowed us to examine extended 285 haplotypes surrounding drug resistance genes, revealing rapid breakdown of haplotypes in the 286 population and different signals of selection in East vs. West DRC.

287

288 We previously investigated population structure using MIPs targeting 20 microsatellites in the DRC¹⁰, 289 failing to detect a strong signal of population structure based upon these markers. Here we leveraged 290 the same 552 samples as the previous study, plus additional samples from the DRC and neighboring 291 countries, to identify clear structure with an improved SNP-based genotyping method. Our ability to 292 detect population structure in the present study is likely due to several factors. First, the new SNP panel 293 contains nearly two orders of magnitude more markers than the previous panel. While this new SNP 294 MIP panel expanded the number of loci interrogated, we have yet to achieve the full potential of MIPs. 295 Specifically, massively increased, multiplexed probe sets that target additional portions of the genome 296 are feasible. MIPs have now been used in human studies to detect as many as 55,000 markers in a 297 single reaction²². Second, a large number of genome-wide SNPs in this study were chosen based on 298 high F_{ST} values in publicly available samples from surrounding countries. This increases our power to 299 detect geographic differentiation, but comes at the cost of not being able to comment on the relative 300 importance of geography vs. drug resistance, which would require random genetic sampling or 301 alternatively whole genomes. Similarly, we should be cautious when interpreting spatial clines in 302 population structure from our data, as we may have greater power to detect structure along some axes

than others due to the unequal distribution of surrounding countries in publicly available samples,
although in general we have good representation in both the East-West and North-South directions.

306 The flexible nature of MIP panels allows for multiplex detection of SNPs associated with drug 307 resistance in any known or putative resistance loci for which they are designed. This allowed for a more 308 detailed evaluation of molecular markers associated with antimalarial resistance than has previously 309 been possible in the DRC. To date, studies of antimalarial resistance markers in the DRC have 310 focused primarily on *pfcrt* (K76T), *dhfr* (N51I, C59R, S108N, I164L), *dhps* (I431V, S436A, A437G, K540E, A581G, A613S), *pfmdr* (N86Y, F184Y, D1246Y), and a few *kelch* mutations^{23–29}. The data 311 312 suggests that mutations associated with artemisinin resistance remained absent in the country as of 313 2014. The World Health Organization identified 9 mutations within the K13 propeller region that are 314 validated in terms of their clinical phenotype of artemisinin resistance, and a further 11 mutations that 315 are candidates associated with the phenotype of delayed clearance.³⁰ We identified 14 mutations within 316 the K13 gene (Supplemental Table 2), although none of these correspond to validated or candidate 317 artemisinin resistance mutations.

318

319 Beyond looking at mutations within drug resistance genes, differences in extended haplotypes around drug resistance genes have been used to understand evolution and spread³¹. Though not originally 320 321 designed for this purpose, the genome wide MIP panel can be leveraged for conducting similar 322 analyses. For example, the differences in CVIET EHH between the West and East suggests that the 323 CVIET haplotype in the West has potentially been more recently introduced, has experienced less 324 breakdown through recombination, or has undergone stronger recent positive selection as compared to 325 the East. Redesign of the selected targets with denser sampling around known drug resistance genes 326 will allow for more robust assessment of these selected regions.

327

328 DRC's location in central Africa and the enormous number of malaria cases in the country means that 329 malaria control in Africa likely depends on improving our understanding on Congolese malaria. This 330 represents the largest study of falciparum population genetics in the DRC and, unlike other large 331 population genetic studies of malaria in Africa, leverages a nationally representative sampling 332 approach. Thus, this study provides the first data on fine-scale genetic structure of parasites at a 333 national scale in Africa, and provides a baseline that can be used to study how implementation 334 programs impact parasite populations in the region. The newly implemented MIP platform represents a 335 highly scalable and cost-effective means of providing genome-wide genetic data, relative to whole

- 336 genome sequencing ¹⁰. The highly flexible nature of the platform allows it to be rapidly scaled in terms
- 337 of targets and samples leading it to be applicable across malaria endemic countries.

338 METHODS

339

340 Study Populations: Chelex-extracted DNA from dried blood spots, collected as part of the 2013-2014 341 DRC Demographic Health Survey (DHS), was tested using quantitative real-time PCR as described 342 previously^{32,33}. Previously published DRC samples¹⁰ were included (n=589), and used to set a Ct 343 threshold of <30 which was applied to the remaining DRC samples (n=1450), resulting in a total of 2039 344 DRC samples sent for sequencing. These samples represented 369 of the overall 539 DHS clusters. In 345 addition, dried blood spot samples from 4 further counties were used: Ghana (n=194), Tanzania 346 (n=120), Uganda (n=63) and Zambia (n=121). Samples from Ghana were collected in 2014 from 347 symptomatic RDT and/or microscopy positive individuals presenting at health care facilities in Begoro (n=94) and Cape Coast (n=98)³⁴. Samples from Tanzania were collected in 2015 from symptomatic 348 RDT-positive patients of all ages at Kharumwa Health Center in Northwest Tanzania³⁵. Samples from 349 350 Uganda were collected in 2013 from RDT-positive symptomatic patients at Kanungu in Southwest 351 Uganda³⁶. Finally, samples from Zambia were collected in 2013 from RDT positive individuals from a 352 community survey of all ages in Nchelenge District in northeast Zambia on the border with the DRC. All 353 non-DRC samples were Chelex extracted, except for the Ghanaian samples which were extracted 354 using QiaQuick per protocol (Qiagen, Hilden, Germany).

355

356 **MIP Design:** We used two distinct MIP panels - a genome-wide panel designed to capture overall 357 levels of differentiation and relatedness, and a drug resistance panel designed to target polymorphic 358 sites known to be associated with antimalarial resistance. The drug resistance MIP panel has been 359 described previously¹⁰. When selecting targets for the genome-wide panel, we used the publicly 360 available P. falciparum whole genome sequences provided by the Pf3k and P. falciparum Community 361 projects from the MalariaGEN Consortium. This consisted of sample sets from Cameroon (n=134). 362 DRC (n=285), Kenya (n=52), Malawi (n=369), Nigeria (n=5), Tanzania (n=66) and Uganda (n=12) 363 (Supplemental Table 1). The genomic sequence from these samples underwent alignment, variant 364 calling, and variant-filtering following the Pf3k strategy consistent with the Genome Analysis Toolkit (GATK) Best Practices with minor modifications^{37–40}. Full details of the bioinformatic pipeline used in 365 366 MIP design are given in the Supplemental Text. Samples from Nigeria and Uganda were dropped 367 after variant calling due to small sample sizes, and the final filtered sequences were used to calculate Weir and Cochran's F_{ST}⁴¹ with respect to country for each biallelic locus. The 1,000 loci with the highest 368 369 F_{ST} values were considered for MIP design as phylogeographically informative loci. Of these 1,000 370 potential loci, 739 were identified as regions that were suitable for MIP-probe design. Separately, from 371 the combined SNP file, we identified 1595 loci that had a minor-allele frequency greater than 5%, had

an F_{ST} value between 0.005 and 0.2, and were annotated by SNPEff as functionally silent mutations.
These loci were identified as putatively neutral SNPs, and 1151 were found to be suitable for MIP
design. The distribution of MIPs is shown in **Supplemental Figure 13** and MIP sequences and targets
are shown in **Supplemental Table 4**.

376

377 Capture and Sequencing: In addition to patient samples, control samples were known mixtures of 4 378 strains of genomic DNA from malaria at the following ratios: 67% 3D7 (MRA-102, BEI Resources, 379 Manasas, VA), 14% HB3 (MRA-155), 13% 7G8 (MRA-154) and 6% DD2 (MRA-156). They were also 380 represented at two different parasite densities (29 and 467 parasites/µI). MIP capture and sequencing 381 library preparation were carried out as previously described¹⁰. Drug resistance libraries were 382 sequenced on Illumina MiSeg instrument using 250 bp paired end sequencing with dual indexing using 383 MiSeg Reagent Kit v2. Genome-wide libraries were sequenced on Illumina Nextseg 500 instrument 384 using 150 bp paired end sequencing with dual indexing using Nextseg 500/550 Mid-output Kit v2. 385 Sequencing reads have been deposited into the NCBI SRA (Accession numbers: pending).

386

Variant Calling and filtering: Variant calling was performed as described previously¹⁰. Within each sample, variants were dropped if they had a Phred-scaled quality score of <20. Across samples, variant sites were dropped if they were observed only in one sample, or if they had a total UMI count of less than 5 across all samples. This data set was considered the final raw data used for additional filtering.

392 Additional filters were applied to both genome-wide and drug resistance datasets prior to carrying out 393 analysis. Sites were restricted to SNPs, and in the case of the genome-wide panel these were filtered 394 to the pre-designed biallelic target SNP sites. Any variant that was represented by a single UMI in a 395 sample, or that had a within-sample allele frequency (WSAF = UMI count/coverage) less than 1%, was 396 eliminated. Any site that was invariant across the entire dataset after this procedure was dropped. 397 Samples were assessed for quality in terms of the proportion of low-coverage sites, where low-398 coverage was defined as fewer than 10 supporting UMIs. Samples with >50% low-coverage loci were 399 dropped. Variant sites were then assessed by the same means in terms of the proportion of low-400 coverage samples, and sites with >50% low-coverage samples were dropped. Samples were then 401 combined with metadata, including geographic information, and were only retained if there were at least 402 10 samples in a given country. This resulted in dropping Tanzanian samples from the drug resistance 403 dataset, but no other countries were dropped. Post-filtering, genome-wide data consisted of 1382 404 samples (DRC = 1111, Ghana = 114, Tanzania = 30, Uganda = 45, Zambia = 82) and 1079 loci, and

drug resistance data consisted of 674 samples (DRC = 557, Ghana = 29, Uganda = 43, Zambia = 45)
and 1000 loci.

407

408 Complexity of Infection: We applied THE REAL McCOIL categorical method to the SNP genotyped
 409 samples to estimate the COI of each individual¹³. Details of the analysis are in the Supplementary
 410 Text.

411

412 **Analysis of population structure:** WSAFs were calculated for all genome-wide SNPs, with missing 413 values imputed as the mean per locus. Principal component analysis (PCA) was carried out on WSAFs 414 using the *prcomp* function in R version 3.5.1. The relative contribution of each locus was calculated 415 from the loading values as $|l_i| / \sum_{i=1}^{L} |l_i|$, where $|l_i|$ is the absolute value of the loading at locus *i*, and 416 *L* is the total number of loci. PCA results were explored in a spatial context by taking the mean of the 417 raw principal component values over all samples in a given DHS cluster, and plotting this against the 418 geoposition of the cluster.

419

420 Identity by descent analysis: Pairwise identity by descent (IBD) was calculated between all samples from the genome-wide SNPs. We used Malécot's⁴² definition of f as the probability of identity by 421 422 descent, where f_{uv} can be defined as the probability of a randomly chosen locus being IBD between 423 samples u and v. At locus i, let A denote the reference allele, which occurs at population allele frequency p_i , and let a denote the non-reference allele, which occurs at population allele frequency 424 $q_i = 1 - p_i$. Assuming that both samples u and v are monoclonal, let X_{ui} denote the observed allele at 425 426 locus *i* in sample *u*, and equivalently let X_{vi} denote the observed allele in sample *v*. Then the 427 probabilities of all possible observed allele combinations between the two samples can be written:

428

429
$$Pr(X_{ui} = A, X_{vi} = A | f_{uv}) = f_{uv}p_i + (1 - f_{uv})p_i^2$$
430
$$Pr(X_{ui} = A, X_{vi} = a | f_{uv}) = (1 - f_{uv})p_iq_i$$
431
$$Pr(X_{ui} = a, X_{vi} = A | f_{uv}) = (1 - f_{uv})p_iq_i$$
432
$$Pr(X_{ui} = a, X_{vi} = a | f_{uv}) = f_{uv}q_i + (1 - f_{uv})q_i^2$$
432

433

from which we can calculate the likelihood of a given value of f_{uv} over all loci as:

435

436 $L(f_{uv} | X_u, X_v) = \prod_{i=1}^{L} Pr(X_{ui}, X_{vi} | f_{uv}).$ (eq2)

437

In practice, population allele frequencies (p_i) were calculated using the mean WSAF for that locus over all samples. Samples were then coerced to monoclonal by calling the dominant allele at every locus.

- 440 The likelihood was evaluated using **eq2** in log-space for a range of values of f_{uv} distributed between 0
- and 1 in equal increments of 0.02. The maximum likelihood estimate $\hat{f}_{uv} = argmax_f L(f | X_u, X_v)$ was
- 442 calculated between all sample pairs. Hereafter the terms "IBD" and \hat{f}_{uv} are used interchangeably.
- 443

Mean IBD was calculated within and between DHS clusters, and compared using a two-sample t-test.
Sample pairs were also binned into groups based on geographic separation (great circle distance) in
100km bins, with an additional bin at distance 0km to capture within-cluster comparisons. Mean and
95% confidence intervals of IBD ware calculated for each group. Finally, sample pairs with IBD>0.9
were identified, and explored in terms of their WSAFs and their spatial distribution.

449

Estimating mutation prevalence from drug resistance panel: Given previous findings of an East-West divide in molecular markers of antimalarial resistance in the DRC^{8,9}, all samples in the DRC were divided by geographically-weighted K-means clustering into two populations. The prevalence of every mutation identified by the drug resistance MIP panel was then calculated in East and West DRC, as well as at the country level. Prevalences in each DHS cluster were used to produce smooth prevalence maps using PrevMap version 1.4.2 in R⁴³, using the method described in Aydemir et. al. (2018)¹⁰.

456

457 Analysis of monoclonal haplotypes: Results of the previous COI analysis on the genome-wide SNPs 458 with THE REAL McCOIL were used to identify samples that were monoclonal with a high degree of 459 confidence. Samples were defined as monoclonal if the upper 95% credible interval did not include any 460 COI greater than one. This resulted in 408 monoclonal samples, of which 143 overlapped with the drug 461 resistance MIP dataset and therefore could be used to explore the joint distribution of mutations in drug 462 resistance genes. 107 of these were from DRC. Analysis focussed on the *dhps* and *crt* genes. Raw combinations of mutations were visualized using the UpSet package in R²¹, and the spatial distribution 463 464 of haplotypes was explored by plotting these same mutant combinations against DHS cluster 465 geoposition.

466

Extended haplotype homozygosity analysis: In order to improve our power to detect hard-sweeps
and capture patterns of linkage-disequilibrium with EHH statistics among putative drug resistance
SNPs, we combined the genome-wide and the drug resistance filtered biallelic SNPs into a single
dataset. Details of this analysis are described in the Supplemental Text.

471

- 472 All associated EHH calculations were carried out using the R-package rehh, and were truncated when
- 473 fewer than two haplotypes were present or the EHH statistic fell below 0.05^{44,45}. In addition, we allowed
- 474 EHH integration calculations to be made without respect to "borders," which were frequent due to the
- 475 MIP-probe design. Although this would result in an inflated integration statistic if the EHH statistic had
- 476 not yet reached 0 within the region of investigation, this problem was mitigated by only comparing
- 477 between subpopulations, and not between loci. EHH decay, bifurcation plots, and haplotype plots were
- 478 adapted from the rehh package objects and modified using $ggplot^{46}$.

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487

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490

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- 498
- 499 Competing Interests: None

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